

Extended abstract

Introduction

Mantle xenoliths exhumed from the Neogene-Quaternary alkaline volcanism in the Afro-Arabian domain provide a powerful tool to unravel the complex mantle dynamics in a region that was interested by plume-related Oligocene Continental Flood Basalt (CFB) magmatism (Hofmann et al., 1997; Pik et al., 1998; Beccaluva et al., 2009) and rifting processes (Natali et al., 2011; 2013), from continental break-up to oceanization.

The influence of the plume has been gradually vanishing over time, as suggested by seismic images of the Afar region, where transition from continental rifting to incipient oceanization has been studied in detail (Bastow et al., 2011; Hammond et al., 2013).

In the Afro-Arabian region, mantle xenoliths included in Neogene-Quaternary alkaline volcanics are found in two main occurrences: i) within the Northern Ethiopian-Yemen plateau area (Beccaluva et al., 2011) and ii) outside this area, neighbouring the rift structures which radiate from the Afar triple junction along the Red sea-Gulf of Aden Arabian margins (Henjes-Kunst et al., 1990; Blusztajn et al., 1995; Chazot et al., 1996; Baker et al., 1998) and along the Main Ethiopian and Kenya-Tanzania rifts southward (Kaesler et al., 2006; Aulbach et al., 2011; Beccaluva et al., 2011; Bianchini et al., submitted).

While mantle xenoliths related to the CFB area show evidence of pervasive refertilization by Afar plume melts (Beccaluva et al., 2011), other mantle xenoliths collected outside this area, along the Arabian margin and the East Africa Rifts, display more complex history of depletion and enrichment events (Baker et al., 1998; Reisberg et al., 2004; Aulbach et al., 2011; Bianchini et al., submitted). In this paper we present new bulk rock and mineral major and trace element data as well as Nd-Hf-Pb-He isotope systematics from an exhaustive sampling (more than 60 samples) from Bir Ali (Yemen), which is a xenolith occurrence located in the southern Arabian margin, ca. 500 km eastward of CFB from the Ethiopian-Yemeni plateau. The aim is to define: i) the real composition, on a statistical basis, of the lithospheric mantle section underlying the southern Arabian margin; ii) its secular variations in terms of depletion and enrichment processes; iii) the geochemical and isotopic signature of the metasomatic events in relation to the Afar plume influence.

Methods

Mantle xenoliths studied in this work have been collected from pyroclastic material of the Bir Ali diatreme, belonging to the Pliocene-Quaternary Balhaf Bir Ali alkaline volcanic province (Mallick et al., 1990), which is located in the central southern part of Yemen (Fig. 1). 62 ultramafic xenoliths (up to 20 cm in size) unaffected by host basalt infiltration were selected for a detailed investigation. Samples were sliced and the freshest portions (weighting between 15 and 30 g) were crushed and then powdered in an agate mill. X-ray fluorescence (XRF) major and trace elements (Ni, Co, Cr, V and Sr) were analysed on powder pellets, using a wavelength-dispersive automated ARL Advant 'X spectrometer at the Department of Earth Sciences of the Ferrara University. Accuracy and precision for major elements are

estimated as better than 3% for Si, Ti, Fe, Ca, and K, and 7% for Mg, Al, Mn, Na; for trace elements (above 10 ppm) they are better than 10%. REE, Sc, Y, Zr, Hf, Nb, Ta, Th, and U were analysed (after HF-HNO₃ dissolution of rock powders in teflon beakers) by inductively coupled mass spectrometry (ICP-MS) at the Department of Physics and Earth Sciences of the Ferrara University, using an X Series Thermo-Scientific spectrometer. Accuracy and precision, based on the replicated analyses of samples and standards, are estimated as better than 10% for all elements well above the detection limit. Mineral compositions were obtained at the CNR-IGG Institute of Padova with a Cameca SX-50 electron microprobe (fitted with four wavelength dispersive spectrometers) at an accelerating voltage of 15 kV and specimen current of 15 nA, using natural silicates and oxides as standards. Trace element analyses on pyroxenes were carried out at the CNR-IGG of Pavia by LAM ICP-MS, using an Elan DRC-e mass spectrometer coupled with a Q-switched Nd:YAG laser source (Quantel Brilliant). The spot diameter was typically 50 μ m, and the CaO content was used as internal standard. Precision and accuracy, better than 10% for concentrations at ppm level, were assessed by repeated analyses of NIST SRM 612 and BCR-2 standards.

For the Nd-Hf-Pb isotopic analysis hand-picked clinopyroxene separates (weight of 100-300 mg) and bulk rock powders (weight of 300-600 mg) have been preliminarily prepared in laboratories of the Department of Earth Sciences at the University of New Hampshire. Samples were leached in hot (~120°C) 6 N HCl to remove any surface contamination (cf. Wittig et al., 2006), following techniques outlined in Blichert-Toft (2001) and Blichert-Toft and Albarède (2009). The resulting residues were subsequently digested in a mixture of concentrated HF-HNO₃. Lutetium (Lu) and hafnium (Hf) were separated as described by Blichert-Toft et al. (1997) and the Light Rare Earth Element (LREE) fraction recovered from the Lu-Hf separation protocol were further processed to separate and concentrate neodymium (Nd). Isotopic measurements were carried out by the Nu Plasma HR multi-collector inductively coupled plasma mass spectrometer (MC-ICP-MS) at the Ecole Normale Supérieure in Lyon. Hafnium and Nd isotope analyses of samples were run in alternation with JMC-475 Hf and “Rennes” in-house Nd (courtesy C. Chauvel) standards, respectively and were normalized for mass fractionation relative to, respectively, $^{179}\text{Hf}/^{177}\text{Hf} = 0.7325$ and $^{146}\text{Nd}/^{144}\text{Nd} = 0.7219$ using an exponential law.

The 100-ppb JMC 475 Hf standard, run throughout the analytical session (n=16) to monitor instrument performance, yielded $^{176}\text{Hf}/^{177}\text{Hf} = 0.282155$ (with external $2\sigma = 0.000010$).

Lead was separated using techniques based on those described in Bryce and DePaolo (2004), and Pb isotopic measurements were carried out on the Nu Plasma HR multi-collector inductively coupled plasma mass spectrometer (MC-ICP-MS) at the Ecole Normale Supérieure in Lyon. For Pb, mass fractionation was corrected via thallium normalization as described in White et al. (2000), and ratios were additionally adjusted for drift using the standard bracketing technique outlined in Albarède et al. (2004) using NIST SRM values reported in Eisele et al. (2003). Four NIST SRM 981 run as “blind” samples amongst the

seventeen bracketing standards analyzed yielded averages (with 2σ external precision) of $^{208}\text{Pb}/^{204}\text{Pb} = 36.7271$ (0.0019), $^{207}\text{Pb}/^{204}\text{Pb} = 15.4978$ (0.0009) and $^{206}\text{Pb}/^{204}\text{Pb} = 16.9408$ (0.0012).

Helium was extracted from hand picked olivine crystals by in vacuo crushing and analysed at the laboratories of the Oregon University (US) using the procedure described by Graham et al. (1998).

Petrography and geochemistry

The composition in terms of the Ol-Opx-Cpx normative classification (following Niu et al., 1997) shows that mantle xenoliths may be classified as spinel (Sp)-peridotites (35 lherzolites, 10 dunites and 6 harzburgites) and Sp-pyroxenites s.l. (5 ol-websterites and 5 websterites). Classification has been crosschecked by mass-balance calculation between bulk rock and mineral chemistry of the constituent phases, showing a good agreement with the normative method. Sp-peridotites are mainly protogranular in texture, only locally porphyroclastic and rarely equigranular. **Olivine** is generally medium to coarse grained (up to 2 mm) and moderately kink-banded, varying in composition from Fo 87.3 to Fo 92.0 in lherzolites, and from Fo 89.1 to Fo 92.9 in dunites. **Orthopyroxene** varies in size from medium to small grained, commonly showing cpx exsolution *lamellae*; it ranges in composition from En 88.9 to En 92.5. **Clinopyroxene** is generally small grained and often interstitial, displaying opx exsolution *lamellae* in the relatively largest crystals; its composition varies in the range En 49.1-52.9 Fs 0.5-3.0 Wo 45.4-48.7. A peculiar clinopyroxene composition is recorded in dunite BA33 (En 43.5-45.6 Fs 0.3-0.7 Wo 53.7-56.1) which is characterized by distinctive high CaO and TiO₂ contents, suggesting interaction with magmatic melts. Dark brown **Spinel** is scarce, with lobate shape showing the following compositional ranges: mg# 76-81 cr# 11-27 in lherzolites, and mg# 71-79 cr# 17-38 in dunites. Interaction with metasomatising agents are evidenced by secondary minerals overimposed on the primary parageneses, particularly widespread in harzburgites and dunites. Reaction textures include “spongy” borders in clinopyroxene, opacized rims around spinel often containing fine-grained aggregates of secondary olivine and clinopyroxene as well as patches containing brownish to yellowish glass, rare plagioclase (An 52-70) and rare disseminated pargasitic amphibole (mg# 88.8-88.9).

Sp-pyroxenites are represented either as individual xenoliths or as discrete domain texturally equilibrated inside peridotites. Like peridotites, pyroxenites exhibit prevalent protogranular texture, in place turning to porphyroclastic and equigranular, although with different composition of constituent minerals with respect to peridotites. In both websterites and Ol-websterites pyroxenes show mutual exsolution *lamellae*. **Clinopyroxene** is the dominant mineral phase, varying in composition between En 47.3 Fs 4.7 Wo 42.9 and En 51.5 Fs 7.0 Wo 47.6. **Orthopyroxene** composition is in the range En 83.8-85.9 **Olivine** is medium grained, weakly deformed and its composition varies in the range of Fo 82.2-83.9. **Spinel** is brown to dark green in color with mg# 49-61 and cr# 18-33. These compositions

conform to those reported for analogous Bir Ali xenoliths by Ali and Arai (2007). Reaction textures similar to those observed in peridotites, although less evident, are also present.

Thermo-barometric estimates for peridotites, based on the Brey and Kohler, (1990) and Kohler and Brey, (1990) algorithms, indicate that Bir Ali mantle xenoliths equilibrated in a range of T(temperature) 900-1100°C and P (Pressure) 9-20 Kbar. These P-T conditions are in agreement with petrographic observations, indicating the ubiquitous stability of spinel in both peridotites and pyroxenites. These estimates also conform with those reported by Ali and Arai (2007) for Bir Ali, Stern & Johnson (2010) for the Arabian Peninsula, and by Conticelli et al. (1999) for Southern Ethiopia.

Peridotite xenoliths plot along the melting depletion trends with a continuous SiO_2 - CaO - Al_2O_3 and TiO_2 decrease from fertile lherzolites approaching the Primitive Mantle (PM) to extremely depleted dunites. Coherently, compatible elements such as Ni, are positively correlated with MgO from lherzolites to dunites as expected for mantle depletion processes by increasing extraction of basic melts. The restitic nature of these dunites after severe partial melting events is favored with respect to a replacive origin, such as that proposed for “dunite channels” in abyssal and ophiolite mantle peridotites (Sühr, 1999; Bernstein et al., 2006; Piccardo et al., 2007; Abily and Ceuleneer, 2013); this is supported by the lack of any chemical, modal and mineralogical compositional gap within the peridotite depletion trend, as well as the non-oceanic setting of the mantle section beneath Bir Ali.

In the same diagrams, pyroxenites are remarkably displaced from PM indicating that significant chemical components such as Al_2O_3 , CaO , TiO_2 , K_2O variably enriched the pristine peridotite mantle section, possibly due to infiltration and interaction with basaltic melts (Fig. 3). The resulting compositions are perfectly comparable with those of pyroxenite mantle xenoliths from other occurrences of the Arabian peninsula (Stern and Johnson, 2010, and references therein), and conform to most pyroxenites occurring in ultramafic massifs, which are generally considered products of cumulus crystallization of basic melts intruding and interacting with the mantle peridotite (Downes, 2007). By contrast, Bir Ali pyroxenite xenoliths show remarkable differences with respect to those from the Northern Ethiopian plateau area (Injibara and Dedessa), which are comparatively enriched in orthopyroxene (Beccaluva et al., 2011).

Middle (M) and Heavy (H)-REE distribution of Bulk rock Chondrite (Ch)-normalized Rare Earth Elements (REE) of Bir Ali xenoliths is positively correlated with the modal clinopyroxene content, as usually observed in mantle rocks (Coltorti et al., 1999; Beccaluva et al., 2001, 2007; 2008; 2011). Accordingly, the M- and H-REE contents range from 0.4 to 1.9 times chondrite in lherzolites, from 0.3 to 0.5 in harzburgites and from 0.2 to 0.6 in most dunites. A notable exception is represented by three dunites (characterized by abundant reaction textures and presence of glass and amphibole) which show M- and H-REE contents ranging from 0.7 to 1.4 times chondrite.

The enrichment in the most incompatible elements (e.g. Light (L) REE) is indicated by the La_N/Yb_N ratio ranging from 0.3 to 2.0 in lherzolites, 0.4-2.6 in harzburgites and 0.7-4.1 in dunites, with the highest ratios corresponding to samples showing the most widespread

reaction textures. Therefore, the $\text{La}_\text{N}/\text{Yb}_\text{N}$ ratio in peridotites may indicate that the interaction with metasomatic agents were slight or negligible in most lherzolites but much more effective in olivine-rich lithologies such as harzburgites and particularly dunites, as expected by experimental results on peridotite permeability (Toramaru and Fuji, 1986).

The reaction domains including newly formed mineral phases and glass indicate that interactions of peridotite matrix with metasomatic agents were relatively recent and did not attained textural re-equilibration.

Pyroxenites show HREE abundances ranging between 3.5 and 12.2 times chondrite, where the absolute REE concentrations increase from Ol-websterite to websterite. They display slightly positively fractionated REE patterns with $\text{La}_\text{N}/\text{Yb}_\text{N}$ increasing from 1.9 to 3.0, in broad relation with the abundance of modal clinopyroxene. These REE patterns together with major elements and mineralogical composition indicate that pyroxenites may represent mantle section remarkably refertilized by basic melts with tholeiitic-transitional affinity at variable melt/matrix ratio. It is important to note that these processes were old enough to permit a textural re-equilibration between peridotitic and pyroxenitic mantle domains.

Ch-normalized REE distribution of **clinopyroxene** from Bir Ali mantle peridotites is reported in Fig. 5. Relatively homogeneous compositions are recorded within each sample, with patterns generally higher and subparallel to that of the relative bulk rock. M- to H-REE flat patterns are recorded in all clinopyroxenes ranging from 5.3 to 14.9 times chondrite. LREE distribution is highly variable with $\text{La}_\text{N}/\text{Yb}_\text{N}$ down to 0.2 for lherzolite unaffected by metasomatism, increasing up to 3.4 for those affected by metasomatic enrichment. As expected, clinopyroxenes in dunites record the most effective LREE metasomatic enrichment with $\text{La}_\text{N}/\text{Yb}_\text{N}$ up to 3.9.

The amphibole REE patterns (in dunite BA8) is characterized by the M-HREE distribution ranges from 23.4 to 24.7 times chondrite and $\text{La}_\text{N}/\text{Yb}_\text{N}$ in the range 4.1-5.5. The related patterns mimic those of clinopyroxene from the same sample, suggesting that the two phases were reaction products of the same metasomatic agent. The REE distribution of glass (in dunite BA33) is characterized by HREE in the range of 18.6-21.9 times chondrite and positive fractionation, with $\text{La}_\text{N}/\text{Yb}_\text{N}$ averaging around 7.0. The above data coherently suggest that these new phases and the related geochemical enrichments were induced by alkali-silicatic metasomatic agents whose effects are evidenced by modal and textural disequilibrium. To constrain the nature of these metasomatising agents, incompatible element modelling has been performed based on the compositions of the most enriched clinopyroxene and pargasitic amphibole using partition coefficients (K_d) mineral/alkaline basic melt from Zack & Brunn (1998) and Dalpe & Baker (1994), respectively. The calculated metasomatising agents are in good agreement with the incompatible element distribution of basic alkaline lavas from Cenozoic volcanic districts of the African plate, whereas they do not fit with carbonatitic melts. Comparison at regional scale shows that metasomatic agents inferred for Bir Ali conform to those calculated for other mantle xenoliths occurrences of the Afro-Arabian domain, invariably characterized by alkali-silicate nature (Baker et al., 1998;

Beccaluva et al., 2007; Beccaluva et al., 2008; Beccaluva et al., 2011; Natali et al., 2013b). By contrast, mantle xenolith occurrences within the Northern Ethiopian Plateau area exhibit the interaction with subalkaline metasomatic agents, similar to CFB related to the Afar Plume activity (Beccaluva et al., 2011).

Hf-Nd-Pb-He isotopes from Bir Ali xenoliths

Hf-Nd-Pb isotopic analyses were carried out on highly selected (hand-picked) clinopyroxene separates, and subordinately bulk rock powders, whereas He isotopic ratio was measured on olivine crystals. Results are discussed on the basis of the notional depleted (DM), high U/Pb (HIMU) and enriched (EM1, EM2) mantle components which refer to the different types of Ocean Island Basalts (OIB) and Mid Ocean Ridge Basalts (MORB) (Zindler and Hart, 1986; Carlson, 1995; Hofmann, 1997; Stracke et al., 2005).

The general distribution of Bir Ali peridotite xenoliths in terms of $^{143}\text{Nd}/^{144}\text{Nd}$ - $^{176}\text{Hf}/^{177}\text{Hf}$ span from the conventional Depleted Mantle (DM) signature or even more depleted compositions (ϵ_{Nd} up to 30.3 and ϵ_{Hf} up to 64.4) to Enriched Mantle (EM) values (ϵ_{Nd} down to -4.0 and ϵ_{Hf} down to 4.0). The lherzolites are characterized by the most depleted supra-chondritic compositions, whereas harzburgites and dunites show enriched compositions coherent with the observed incompatible element distribution. This isotopic distribution confirm that lherzolites and harzburgites/dunites represent the least and most metasomatized peridotite mantle domains, respectively.

Model ages have been calculated on the basis of $^{176}\text{Hf}/^{177}\text{Hf}$ and $^{143}\text{Nd}/^{144}\text{Nd}$ respect to both CHUR and DM, for the most LREE depleted clinopyroxene of lherzolites BA37 and BA43. Hafnium model ages are in the range 2040-1800 and 1670-1115 Ma, respect to CHUR and DM. Nd model ages are comparatively younger approaching 1300 (CHUR) and 1050 (DM) Ma. The age discrepancy between Hf and Nd model ages is attributable to the different susceptibility of the Lu-Hf and Sm-Nd systems to the metasomatic agents, which are characterized by a strong LREE (e.g. Nd) enrichment.

Therefore, from these estimates we may conclude that significant partial melting events took place during Proterozoic ages, and were followed by metasomatic interactions which variously perturbed the relative isotopic systems.

The Pb isotope systematic generally conforms to the above scenario with many lherzolites plotting close to DM, whereas harzburgites, dunites and websterites generally cluster near the EM components. In particular, the same lherzolites considered for the model ages are those showing unradiogenic Pb isotopic composition, thus confirming that they escaped significant metasomatic effects. More radiogenic Pb compositions are displaced toward the EM2 isotopic end-member. For comparison we report the isotopic compositions of mantle xenoliths from other localities of the Arabian plate (Ataq; Baker et al., 1998; Jordan; Shaw et al., 2007), East Africa (Mega, Southern Ethiopia; author's data; Assab, Eritrea; Teklay et al., 2009; Injibara and Dedessa, Northern Ethiopia; author's data) and from the Saharan Belt (Hoggar, Algeria:

Beccaluva et al., 2007; Gharyan, Lybia: Beccaluva et al., 2008). The resulting distribution show that the depletion and enrichment processes that affected the Afro-Arabian lithospheric domain produced an extreme variability of isotopic signatures, extending sometimes beyond the mantle arrays defined by the conventional mantle isotopic components (DM, EM and HIMU).

A relevant exception is represented by mantle xenoliths located within Northern Ethiopian-Yemeni plateau (Assab, Injibara and Dedessa), showing a relative isotopic homogeneity ($^{176}\text{Hf}/^{177}\text{Hf} = 0.28277\text{-}0.28340$, $^{143}\text{Nd}/^{144}\text{Nd} = 0.51284\text{-}0.51329$, $^{208}\text{Pb}/^{204}\text{Pb} = 37.7\text{-}39.1$, $^{207}\text{Pb}/^{204}\text{Pb} = 15.5\text{-}15.6$, $^{206}\text{Pb}/^{204}\text{Pb} = 17.7\text{-}19.0$), possibly inherited by pervasive interaction of these mantle sections with Afar plume-related CFB which display similar isotopic composition ($^{176}\text{Hf}/^{177}\text{Hf} = 0.28290\text{-}0.28319$, $^{143}\text{Nd}/^{144}\text{Nd} = 0.51284\text{-}0.51307$, Meshesha and Shinjo, 2007 and 2010; $^{208}\text{Pb}/^{204}\text{Pb} = 37.6\text{-}39.1$, $^{207}\text{Pb}/^{204}\text{Pb} = 15.4\text{-}15.7$, $^{206}\text{Pb}/^{204}\text{Pb} = 18.0\text{-}19.3$; author's data and Pik et al., 2006).

The helium isotope composition ($^3\text{He}/^4\text{He}$) of the olivines from Bir Ali varies from 7.5 to 7.8 Ra falling in the range recorded in other occurrences of the East African-Arabian domain (e.g. Mega, Injibara and Dedessa: 6.9-8.9, Beccaluva et al., 2011) and is systematically higher than that measured in peridotite xenoliths from Saharan belts (6.2-6.8; Azrou, Morocco: Natali et al., 2013b; Hoggar: Beccaluva et al., 2007; Gharyan: Beccaluva et al., 2008). A possible explanation is provided by the extreme mobility (and decoupling) of noble gases with respect to other incompatible elements, which may result in a much more extended lithospheric region influenced by the plume. The related diffusive aureola could account for the systematic higher $^3\text{He}/^4\text{He}$ values of East-Africa/Arabia with respect the north African belt.

Conclusions

The extensive sampling of mantle xenoliths from Bir Ali show that spinel-peridotites represent the preponderant part (85%) of the southern Arabian lithosphere, the remaining part (15%) being constituted by spinel-pyroxenites sometimes forming texturally equilibrated composite xenoliths. Peridotites exhibit a compositional variation from lherzolites (up to 14% clinopyroxene) to harzburgites and dunites (cpx down to 2-3%) which delineate a continuous depletion trend attributable multiple and intensive extraction of basic melts. The calculated model ages suggest that these partial melting events occurred at least since Paleo-Proterozoic (2 Ga according to Lu-Hf model age of the most depleted peridotites). Moreover, The Lu-Hf and Sm-Nd error-chrones calculated on the total xenoliths population suggest important magmatic events between 800 and 750 Ma. While the Paleo-Proterozoic ages may indicate the occurrence of extensive partial melting events in the pristine mantle, the Neo-Proterozoic ages may correspond to Pan-African magmatic events that modified the lithospheric mantle, as suggested by Stern & Johnson (2010). In our view, texturally equilibrated pyroxenites, which represent ubiquitous and significant components of mantle xenoliths throughout the

Arabian Peninsula, could be related to these Neo-Proterozoic events that extensively rejuvenated the Arabian lithospheric mantle.

On the other hand, the observed disequilibrium textures and formation of new phases (glass, clinopyroxene and amphibole) in parallel with incompatible elements and isotopic enrichments could be derived by interactions with much younger and recent metasomatic agents that variously affected the studied lithospheric section. These metasomatic effects, particularly effective in harzburgite and dunite lithologies, indicate that the causative agents were OIB-type alkaline basic melts. Therefore, the Bir Ali mantle section, compared with others from the Afro-Arabian domain, confirms that the general extensional regime radiating from the Afar triple junction and generating the Red Sea-Gulf of Aden-Main Ethiopian Rift system, was accompanied by shallow mantle upwelling events and metasomatic processes dominated by alkaline agents (Henjes-Kunst 1990; Blusztajn et al., 1998; Kaeser et al., 2006; Bedini et al., 2007; Shaw et al., 2007; Aulbach et al., 2011; Beccaluva et al., 2011). A notable exception is represented by mantle xenoliths included in Neogene-Quaternary alkaline volcanics located within the Northern Ethiopian-Yemeni CFB province; in these xenoliths, petrological and geochemical evidences indicate that the causative agents of mantle metasomatism were subalkaline melts closely resembling the tholeiitic magmas related to the Afar plume (Beccaluva et al., 2011 and references therein). We may conclude that the thermo-chemical effects of the Afar plume were essentially confined to the lithospheric sections located within the northern Ethiopian-Yemeni CFB plateau, and were negligible in the neighbouring regions. However, based on the available data on extremely mobile elements such as noble gases (Fig. 8), we emphasize a regional scale anomaly throughout the East-African-Arabian domain showing a systematic higher Helium isotopic composition with respect to other African occurrences (residual plume influence?).